87-140943/20 B05 SOUG-02,10.85 16 2081-365-A

B(10-A9B, 12-G1A, 12-H3)

SOUGO YAKKOU KK

02.10.85-JP-218009 (14.04.87) A61k-31/18 C07c-161 Guanidina ethane thiosulphonic acid chalesteral decreasing agent prepd. by reacting guanidino ethane sulphinic acid with sulphur in presence of base C87-058856

Guanidinoethanethiosulphonic acid of formula [1] is new:

$$CH_{2} - N - C NH_{2}$$

$$CH_{3} - SO_{3}SH$$

$$CH_{4} - SO_{3}SH$$

$$(11)$$

USE/ADVANTAGE

If is useful as cholesterol decreasing agent.

The compound has strong cholesterol decreasing activity and strong HDL-cholesterol increasing activity without toxicity (LDso = 3000 mg/Kg in the rat).

PREPARATION

Cod. [1] is prepared by reacting hypotaurocyamine (guanidinoethanesulphinic acid) with sulphur in the presence

of base.

Caustic alkali such as NaOH, KOH is used as bose. Powdered sulphur is pref. used.

Solvent is prof. an alcohol such as McOH. EtOH or i-PrOII.

ACTIVITY

Test results on male rats allowed to eat normal food, cholesterol food, and cholesterol food with 111 (200 mg/Kg. day) for 2 weeks [total] cholesterol in serum, HDL-cholesterol in serum, HDL-cholesterol (mg/dl)] are: 109.2, 48.51 521.2, 20.5; 283.9, 28.1.

EXAMPLE

Hypotaurocyamine (0.48 mol) was dissolved in 0.28 E(OH (1800 ml) and sulphur (6.3g) were added. The mixture was stirred under retlux until the sulphur completely disappeared and was allowed to stand overnight. Crude crystals were filtered and washed with US; (twice) and EtOH. The crystals were dissolved in hotwater and recrystallized by adding EtOH (2700ml) and cooling. Filtration and washing with other afforded 26.4 g (80.1%) of [1], mp 206-210°C. (4ppW67LDDwgNo0/0).

87-140944/20 TOHYOH STAUFER CHEM

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TOST- 02.10.85

02.10.85-JP-219681 (14.04.87) C07d-205/08 Highly stereoselective synthesis of beta-lactam deriv. - by treating lithium enolate of organic ester with organic imine cpd. in polar solvent

C87-058857

8-Lactam derivs, are synthesized highly selectively by treating lithium enolate of organic ester with organic imine epd. in polar solvent.

The organic imine cpd. may be an imine coordinated with trialkylaluminum. When the cpd. is used as imine, cis prod. may be synthesized with 100% stereoselectivity.

USE/ADVANTAGE

Luctams are formed with high stereoselectivity. Prods. are useful as pharmaceuticals.

EXAMPLE

n-BuLi (15% hexane soln.) (12 m mols.) was added to a soln, of disopropylamine (12 m mots.) in n-hexane (7 ml) with ice-cooling under N2, and resultant mixt, was stirred. n-Hexane was distilled off under reduced press., THF (5 ml) was added to the residue, and the mixt, was cooled to -78°C.

B(7-D1)

B0171

(CH,),CHCH,COOC,H, or CH,CH,COOC,H, (10 m mols) was added within three minutes to the above mixt., and a soln, of C. H, CH=NC. H, (10 m mols) in THF (5 ml) or a soln. of the Imine (10 m mols) and AIR, (see below). (10 mmols) in THF (5 m mols) was added.

The low temp, cooling both was removed and temp, of reaction mixt, was elevated slowly to room temp, over ten hours. The mixt, was then hydrolysed with 1N HCl aq. soln. and prod. was extracted with benzene to give 8-lactam.

Yield of the 8-lactam and results of eis: trans ratio are as follows:

(n) R1 = 1-Pr:

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AIR,	Yield (%) 87	Cis: trans ratio		
None		0 : 100		
AI(CII,),	73	100 : 0		
A1(C2115),	75	100 : 0		
Ali-Bu,	40	100 : 0		

(b) R = Cli;

AIR,	Yield (%)	Cis :	trans i		rntio
None	92	0	:	100	
A1(CH,),	85	100	:	0	
A1(C,11,),	83	100	:	0	
Ali-Bu,	5 2	100	:	0	

(5ppW69EDDwgNo0/0).

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